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(54) Dendritic lysine-based polypeptides for targeted drug delivery Dendritische, lysinhaltige Polypeptide zur gezielten Arzneimittelabreichung Polypeptides dendritiques à base de lysine pour apport ciblé de médicaments

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#### Description

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[0001] The present invention relates to polypeptide compounds which have dendritically linked units formed from amino acids having reactive groups, for instance carboxylic acid or amine groups, in their side chains. Each molecule comprises two dendrons. To at least two of the terminal branches of one of the dendrons there are attached anchor groups, each of which comprises at least one lipophilic group. The terminal units of the at least one other dendron may be unconjugated or may be conjugated to ligons of various types. The dendrons are attached at a core which may include a linear oligo peptide, optionally having pendant sugar molecules.

[0002] Tam et al, in Proc.Nat.Acad.Sci.USA(1988) 85, 5409-5413 describe a compound including a dendritically linked polylysine component, to the focal lysine of which is attached a lipophilic moiety, through a peptide bond to the carboxylic acid group of that lysine unit. To the terminal branches of the dendritic moiety there may be attached peptide antigens to provide an active ingredient for a vaccine having improved antigenicity. WO-A-95/00540 describes dendritic carriers, in which the dendrons include lysine or other di-amino carboxylic acids. The carrier may include a hydrophobic group connected by a linker to the focal group of the dendritic molecule. Where the dendritic carrier is synthesised using solid state peptide synthetic methods, the hydrophobic group is linked following cleavage of the dendrimer from the carrier. Suitable hydrophobic groups are derived from fatty acids or fatty alcohols.

[0003] In WO-A-94/02506, Toth et al describe an improvement of Tam's invention, in which the anchor component is formed from lipophilic amino acids. This allows the compound to be synthesised using conventional solid state peptide synthetic techniques, in the first stages of which the lipophilic amino acids are linked to form, for instance, a three unit linear oligo peptide, a focal lysine unit is joined to the final lipophilic amino acid and the dendritic core moiety is then linked to the two amine groups of the focal lysine unit. The peptide antigens may subsequently be synthesised directly onto the terminal branches of the dendritic core, all the steps being carried out without cleavage of the polypeptide from the solid substrate carrier. The synthetic process used to make Toth et al's product required the use of starting amino acid reagents with the same protecting group blocking the two amine groups of lysine reagents. Consequently during the steps in which the dendritic component is synthesised, the same reagent is added to each of the amine moieties.

[0004] In the product of Toth *et al* it was essential for the lipidic amino acids to be joined directly to one another by peptide bonds, and that a lipidic amino acid can be joined to a carrier substrate so that synthesis involves linkage of that unit to the carrier by a peptide bond and linkage of another lipid amino acid unit to the dendritic moiety by a peptide bond. Consequently solid state peptide synthesis methods can be used to conjugate each of the components of the final product to one another. By contrast, in Tam *et al*, whilst the dendritic polylysine and the peptide antigen can be synthesised using solid state peptide synthetic methods, the polylysine-polyantigen compound must be cleaved from the carrier substrate prior to conjugation to the lipophilic anchor moiety, through the carboxylic acid unit of the focal lysine group. The reagent, from which Tam's lipophilic anchor is synthesised, has only one reactive group.

[0005] A new dendritic compound according to the present invention comprises a core including a local group from which at least two dendrons extend, each dendron comprising dendritically linked amino acid units I

in which  $R^1$  is  $C_{1.6}$ -alkylene and X is selected from the group consisting of -O-, -S-, -NH- and -CO-, and each unit of a dendron may have the same groups  $R^1$  and X, and in which a first dendron had n (where n is 2) levels of dendritically linked amino acid units and  $2^n$  terminal branches, to p (where p is at least 2) of which terminal branches there are linked anchor groups

where Y is selected from -CO-, -NH-, -O- and -S-, provided that at least one of X and Y is -CO-,

R<sup>2</sup> is an organic group containing at least one C<sub>6-24</sub>-alkyl, -alkenyl, or -alkynyl group, R<sup>3</sup> is selected from the group consisting of hydrogen, amine, blocked amine, hydroxyl, C<sub>1-24</sub> alkoxy, thiol, COOH, or an organic group containing at least one C<sub>6-24</sub>-alkyl, alkenyl or -alkynyl group, C<sub>1-6</sub>-alkanoyloxy, or C<sub>1-6</sub>-alkanomido

and in which a second dendron has m (where m is in the range 3-5) levels of dendritically linked amino acid units of the formula I above in which the groups R<sup>1</sup> and X may be the same as or different to one another and the same as or different to those of the amino acid units in the first dendron and 2<sup>m</sup> terminal branches, each of which is either unconjugated and is a group selected from NH<sub>2</sub>, N+H<sub>2</sub>R<sup>11</sup>, in which R<sup>11</sup> is hydrogen or C<sub>1-4</sub> alkyl, COOH, COO-, OH or SH, or is conjugated via the terminal -X-, -NH- or -CO- group to a group R<sup>12</sup> where R<sup>12</sup> is a methylol group, an active ligand or an organic group comprising a sugar moiety.

[0006] In the present invention the focal group of the core is linked through covalent bond to the at least 2 dendrons. The core may include components other than the focal unit, for instance joined to the focal unit by one or more additional covalent bond. Preferably the focal group is an amino acid unit, for instance having the formula I

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in which X and R1 are as defined above.

Preferably X is either -NH- or -CO-. Where it is -CO-, the two dendrons are attached to the two -CO- moieties. Where X is NH, the two dendrons are linked one each to the groups NH. Preferably the focal group is formed from lysine or ornithine.

[0007] The core may comprise units other than the focal group. Such units are preferably peptide linked amino acid based units. Additional core units may function merely as spacers, or may include functional groups such as lipophilic groups, hydrophilic groups or active ligands, for instance targeting groups. The compound may be attached to a resin through the focal unit, for instance via a spacer.

[0008] The present invention is made possible by the use in the synthesis of the dendritic compound of a reagent for forming the focal group which has at least three reactive groups, each of which can be sequentially reacted. Where, in the preferred embodiment of the invention, the focal group is an amino acid unit of the formula I where the group X is -NH-, the reagent from which the focal unit is derived has the two amine groups protected by two different protecting groups which are removable under different conditions. Each amine group can consequently be protected, activated and reacted in sequential series of reaction steps. This allows two different dendrons to be synthesised.

[0009] The present invention includes also a method for synthesising the novel compound in which a focal reagent which has two reactive groups is reacted in a first series of first dendron producing steps as follows:

1. an amino acid reagent of the formula II

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in which R1 and X are as defined above,

R14 is H when X is -O-, -S- or -NH-,

OH when X is -CO-, or

is a protecting group,

R15 is a carboxylic acid protecting group, hydroxyl, or a carboxylic acid activating group,

R16 is H, an amine protecting group or an amine activating group,

provided that at least two of R14, R15 and R16 is other than an activating group and at least one of R14, R15 and R16 is other than a protecting group, is reacted

with the focal reagent, optionally after a step in which the desired reactive group of the focal reagent and/or one of the groups -XR<sup>14</sup>, -COR<sup>15</sup> and -NHR<sup>16</sup> is deprotected and/or activated whereby the reactive groups on the focal reagent reacts with one of the groups R<sup>14</sup>X-, R<sup>15</sup>-CO- and R<sup>16</sup>-NH-;

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2. a second step in which both unreacted groups R14X-, R15CO- and R16NH- of the product of the preceding step are, if necessary, deprotected and/or activated and reacted with at least two equivalents of a trifunctional reagent having the general formula II,

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in which the groups R1, R14, R15 and R16 are as defined in step 1 and are the same or different as in the trifunctional reagent used in step 1;

- 3. One repeat of step 2, using at least four equivalents of trifunctional reagent; and
- 4. an anchor group attachment step in which at least two of the four groups R<sup>14</sup>X-, R<sup>15</sup>CO- and R<sup>16</sup>NH are, if necessary, deprotected and/or activated, and reacted with a lipophilic reagent of the formula III

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in which Y, R<sup>2</sup> and R<sup>3</sup> are as defined above, and R<sup>17</sup> is OH or a carboxylic acid activating group, where Y is -CO- or is H or an amine, hydroxyl or thiol activating group, respectively, where Y is -NH-, -O- or -S-, whereby the said at least two groups react with R<sup>17</sup>Y-to conjugate

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groups to the X, CO- or NH-; and

a second dendron forming series of reaction steps in which

1. the other of the reactive groups of the focal reagent is, in a step separate to step 1 mentioned above, reacted with an amino acid reagent of the formula II

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in which R1 and X are as defined above,

R14 is H when X is -O-, -S- or -NH-,

OH when X is -CO-, or

is a protecting group,

R15 is a carboxylic acid protecting group, hydroxyl, or a carboxylic acid activating group,

R16 is H, an amine protecting group or an amine activating group,

provided that at least two of R<sup>14</sup>, R<sup>15</sup> and R<sup>16</sup> is other than an activating group and at least one of R<sup>14</sup>, R<sup>15</sup> and R<sup>16</sup> is other than a protecting group,

with the focal reagent, optionally after a step in which the desired reactive group of the focal reagent is deprotected and/or activated whereby the other of the reactive groups on the focal reagent reacts with one of the groups R14X-, R15-CO- and R16-NH-;

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2. a second step in which both unreacted groups R<sup>14</sup>X-, R<sup>15</sup>CO- and R<sup>16</sup>NH- of the product of the preceding step are, if necessary, deprotected and/or activated and reacted with at least two equivalents of a trifunctional reagent having the general formula II,

in which the groups  $R^1$ ,  $R^{14}$ ,  $R^{15}$  and  $R^{16}$  are as defined in step 1 and are the same or different as in the trifunctional reagent used in step 1;

3. (m-1) repeats of step 2, using in each case at least 2<sup>(r+1)</sup> equivalents of trifunctional reagent for the r<sup>th</sup> repeat of step 2, until m levels of dendritically linked amino acids have been formed, where m is in the range 3 to 5.

[0010] In a preferred reaction there is a preliminary step of reacting a focal reagent of the formula VI

$$R^{17}-X-R^{1}-CH-NH-R^{18}$$

| VI

in which R<sup>17</sup>, R<sup>18</sup> and R<sup>19</sup> are selected from the same groups as R<sup>14</sup>, R<sup>16</sup> and R<sup>15</sup>, respectively, as defined above, with a substrate having a pendant group which is capable of reacting with one of the groups XR<sup>17</sup>, NHR<sup>18</sup> and COR<sup>19</sup>, optionally after deprotection and/or activation of the said pendant group or said one of the groups of the focal reagent, whereby the focal reagent is bound to the substrate.

[0011] In the process, the series of reactions used to form the dendron having lipophilic moieties  $R^2$  may be carried out before or after the series of reactions to form the other dendron. Thus the reference to the first series of steps and second series of steps does not, unless the context makes it explicit, imply an order of carrying out the said series.

[0012] In the invention, in the lipophilic component -Y-CH( $R^2$ ) $R^3$ ,  $R^2$  is preferably selected from  $C_{6-24}$ -alkyl, - alkenyl or - alkynyl, or is a group IV

in which R4 is a bond or a C1-6-alkylene group,

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 $R^5$  is hydrogen, a  $C_{1-6}$ -alkyl or a  $C_{1-24}$ -alkanoyl group or a group  $CH_2SCH_2CH(OCOR^{20})CH_2OCOR^{20}$ , in which  $R^{20}$  is a  $C_{6-24}$ -alkyl, -alkenyl or -alkynyl group,  $R^6$  is hydrogen or a  $C_{6-2-4}$ -alkyl, -alkenyl group,  $R^6$  is hydrogen or a  $C_{6-2-4}$ -alkyl, -alkoyy, -alkanoyl group,  $R^6$  is hydrogen or a  $C_{6-2-4}$ -alkyl, -alkoyy, -alkanoyl group,  $R^6$ -alkyl, -al

 $\mathsf{R}^6$  is hydrogen or a  $\mathsf{C}_{6\cdot 24}\text{-alkyl}$ , -alkoxy, -alkanoyl or-alkanoyloxy group or  $\mathsf{R}^2$  is a group

$$R^{21}$$
-CO-NH- $R^7$ -O-P-O- $R^8$ - $R^9$  ( $R^{10}$ ) 2

in which  ${\sf R^8.\ R^{21}}$  and  ${\sf R^7}$  are each  ${\sf C_{2-6}}$ -alkylene  ${\sf R^9}$  is glyceryl and

each group  $R^{10}$  is independently selected from  $C_{6-24}$ -alkyl, -alkenyl, -alkynyl, -alkanoyl, -alkenyl or -alkynoyl, provided that  $R^5$  and  $R^6$  cannot both be groups selected from hydrogen, lower alkyl, alkenyl and alkynyl groups.

[0013] Where the group  $\mathbb{R}^2$  is a group of the formula IV, and especially where  $\mathbb{R}^4$  is a bond, the compound is derived from a lipidic amino acid of the formula V

HOOCCH-R<sup>6</sup>

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wherein R6 is as defined above.

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[0014] In the step in which a lipidic amino acid is conjugated to the terminal branches of the first dendron whichever of the COOH and NH<sub>2</sub> group is not desired to react with the terminal branch is generally blocked by an appropriate protecting group.

[0015] It may often be more convenient to use, instead of the lipidic amino acid of the formula V, a monofunctional reagent to provide the anchor moieties, for instance a fatty acid or fatty amine.

[0016] The dendritic compound of the invention may be bound to a solid support, for instance a resin used as the solid peptide synthesis support. Thus the core is joined to the solid support, for instance a resin, through the focal unit, optionally via a spacer, for instance an oligopeptide spacer. The compound may be cleaved from the support prior to use, optionally after having reacted further any of the underivatised terminal branches. Thus the unreacted terminal branches may be in the form of free or protected carboxylic acid, amine, hydroxyl or thiol groups.

[0017] In a preferred aspect of the invention the dendritic compound has several terminal primary amino groups or is in the form of the corresponding ammonium salt. Usually all the terminal groups of the second dendron are amine or ammonium groups.

[0018] In a further preferred embodiment of the product of the invention, at least some of the terminal groups of the second dendron are attached to an organic group comprising a sugar molecule.

[0019] Generally it is preferred for all of the terminal branches of the first dendron to be provided with lipophilic anchor moieties. It is found that two or four such moieties are adequate to provide appropriate levels of lipophilicity to the compound as a whole.

The second dendron has at least three levels of dendritically linked amino acid units, preferably four or, sometimes five levels of dendritically linked amino acid units (that is, m is 3 to 5). Where there are five or more levels of dendritically linked amino acid units, stearic hindrance may prevent full dendritic linkage of groups, for instance further dendritic moieties, to the terminal units. Consequently it is preferred for there to be no more than five, and preferably four, levels of dendritically linked amino acid units.

[0021] As indicated below in the detailed examples, it has been found that the dendrimer of the present invention having four anchor groups being lipidic amino acid units joined to the amine terminal groups of the first dendron, and with free amino groups at each of the terminal groups of 3-, 4- and 5- level dendritically linked amino acid units for the second dendron have reduced toxicity as determined by enythrocyte lysis, as compared to a lipid peptide dendromer as described in our earlier application WO-A-94/02506 comprising a linear oligopeptide anchor moiety of three lipidic amino acids joined to the focal lysine of a dendrimer having the equivalent number of levels of dendritically linked amino acid (lysine) units.

[0022] The compound of the invention has a similar utility to those described in WO-A-94/02506. Thus, to the terminal branches of the second dendron, there may be conjugated peptide antigens, drug moieties, targeting moieties, for instance antibodies or sugar groups, or other groups providing increased levels of hydrophilicity (for instance sugar molecules, polyethylene glycol molecules or ionic moieties).

[0023] The invention is illustrated further in the following examples.

#### MATERIALS AND METHODS

[0024] Polystyrene aminomethylated (PAM) resin, BOC-protected aminoacids from Novabiochem, 2-(1H benzotriazole-lyl)-1,3,3,-tetramethyluronium hexafluorophosphate (HBTU) from Phase Separations Ltd, Trifluoroacetic acid (RFA) from Halocarbon Products Corporation, hydrogen fluoride gas (HF) from BOC, diisopropyl ethyl amine (DIEA) from Fluka and dimethylformaamide (DMF) from Rathburn were all used as received. The protected lipidic aminoacids were synthesised and purified in our laboratory as described in Gibbons, WA, et al (1990) Liebigs Ann.Chem. 1177-1183.

#### Example 1

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[0025] Solid phase peptide synthetic methods were used employing a polyacrylamide resin having a degree of substitution of 0.46 mmol/g resin. The reaction sequence is shown in slow diagram Figure 1 the step involving protection of Boc was performed in 100% trifluoroacetic acid. Couplings of pendant amine groups on the bound compound with carboxylic acid groups of amino acid reagents having protected amine groups was achieved using a three fold excess of HBTU activated Boc-amino acids in dimethylformamide in the presence of disopropylethyl amine. Acidulation of

deprotected Boc group of lipoamino acid was carried out in the presence of disopropylethyl amine. Deprotection of the Fmoc group to form the second dendron was carried out by a suitable system.

[0026] The resin peptide was carefully flow washed before and after each deprotection step. The final product was washed with dichloromethane and dried in air. The peptide was removed from the resin support with a high HF method 2 g resin peptide, 20 ml HF, 1.5 hour at -5°C) to yield the crude peptide which was dissolved in 95% acetic acid solution and lyophilised.

#### Purification

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[0027] Analytical HPLC separation of the synthesised dendrimers was carried out on a 25 cm Vydac C<sub>18</sub> RAC column with 5 µm pore size and 4.6 mm internal diameter. Following standard degassing techniques, particulate matter was removed from HPLC grade acetonitrile and water using membrane filters. Analytical separation was achieved with a solvent gradient beginning with 0% acetonitrile, increasing to 60% acetonitrile at 20 min. maintaining at this concentration for 20 min and decreasing steadily to 0% acetonitrile for 10 min at a constant flow of 1.2 ml min<sup>-1</sup>. For preparative separation a TSK-GEL preparative C<sub>18</sub> column with 10 µm pore size and 2.5 cm internal diameter was used. Separation was achieved with a solvent gradient beginning with 0% acetonitrile, increasing constantly to 18% acetonitrile at 60 min then 60% acetonitrile at 80 min, staying at this concentration for further 30 min and decreasing steadily to 0% acetonitrile for 30 min at a constant flow of 8 ml min<sup>-1</sup>. The gradient was effected by two microprocessor-controlled Gilson 302 single piston pumps. Compounds were detected with a Waters 486 tunable absorbance detector at 214 nm or a Holocrome UV-VIS detector 220 nm. Mass spectra were run on VG Analytical Tofspec instrument, using matrix assisted laser desorption (MALD) ionisation at a wavelength of 337 nm generated by a nitrogen laser.

[0028] Compound synthesis using the general technique mentioned above had the following general structure:

[0029] The compounds synthesised have the values for the number of lipid residues and the number of primary amine groups as well as the molecular weight shown in Table 1.

TABLE	4
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Compound	R	Levels of dendritically linked lys residues in 2nd dendron	No 1° amine groups	MW
1.1	NH <sub>2</sub>	3	8	2500
1.2	Lys(NH <sub>2</sub> ) <sub>2</sub>	4	16	3526
1.3	Lys(Lys(NH <sub>2</sub> ) <sub>2</sub> ) <sub>2</sub>	5	32	5577

#### Comparative Example 1

[0030] Using the same general techniques described above in relation to Example 1, but omitting the Fmoc strategy, compounds having the general formula shown below were produced. Thus the process involved three sequential steps to provide a linear tripeptide of lipoamino acid units bound to the glycine group attached to the resin, followed by a step of adding a Boc Lys (Boc) OH unit to the third lipoamino acid unit, followed by deprotection of both the amine groups of lysine and addition of sequential dendritically linked lysine moieties.

[0031] The methylol compound was synthesised by subjecting the compound having eight free amine groups at the terminal ends to reaction with a suitable reagent. The compounds synthesised are shown in Table 2 below.

T	Ά	В	L	Ε	2

Compound	R'	Levels of dendritically linked Lys (n)	No 1° amine groups	MW
1.4	NH <sub>2</sub>	3	. 8	1590
1.5	Lys(NH <sub>2</sub> ) <sub>2</sub>	4	16	2615
1.6	Lys(Lys(NH <sub>2</sub> ) <sub>2</sub> ) <sub>2</sub>	5	32	4666
1.7	СН₂ОН	3	0	3390

#### Rat Erythrocyte Lysis Studies

[0032] Fresh blood was obtained from rats through cardiac puncture, collected in heparinised tubes and centrifuged at 1,000 g for 15 minutes at 4°C. The supernatant, was discarded, the volume was made up to 10 ml with chilled phosphate buffered saline (PBS). The suspension was centrifuged again and the PBS washing step was repeated twice. Finally, the supernatant was removed and the cell pellet resuspended up to 2% w/v in chilled PBS. 100 µl of samples of compounds 1.1-1.7 of different dilutions were added in flat bottomed Elisa plate. 1% w/v of Triton X 100 was used as the control (100% lysis). 100 µl of erythrocyte suspension was added and incubated for 1h, 5h and 24 hrs. At different time intervals these plates were removed and the suspensions centrifuged. 100 µl of the supernatant was removed and placed into fresh Elisa plate and the absorbance was measured at 545 nm with PBS as blank. The % population lysis was calculated by using the formula

Percentage population lysis = (Absorbance/control (triton) absorbance) 100.

#### Results

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[0033] The toxicity of compounds 1,1-1.7 were compared with linear polylysine of two different molecular weights (34,000 and 1000-4000). Triton X100 was used as positive control. The higher M.W. polylysine had a concentration independent toxicity 35.7% to 54.2% of percent population lysis was observed between the concentrations 1 µg/ml to 30 µg/ml. The lower M.W. polylysine was found to be almost nontoxic.

[0034] Red blood cellsysis studies indicated that compounds 1.4-1.6 were non toxic at the low concentration of 1 µg/ml after 24 hrs where as at higher concentrations (above 20 µg/ml) these compounds were toxic even after one hour incubation. All compounds 1.1-1.6 had concentration dependent toxicity.

[0035] The toxicity studies of compounds 1.1 to 1.3 showed that the toxicity is dependent on the ratio of the lipophilic groups to the number of amino groups attached to the molecule. Compound 1.1, which contained 8 amino groups found to be less toxic than similar compounds having 16 and 32 amino groups (1.2 and 1.3) were less toxic than comparative 1.4 to 1.6. This indicates that although compounds 1.1 to 1.3 were bulkier, the position of attachment of lipo amino acid makes them less toxic.

[0036] Compound 1.7 which contained 3 lipo amino acid chain attached consecutively was non toxic at the concentration of 30 µg/ml up to 5 hours incubation, indicating that the toxicity is not due to the presence of lipo amino acid.

[0037] The results are illustrated graphically in Figures 2 and 3.

Claims

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 A dendritic compound comprising a core including a focal group from which at least two dendrons extend, each dendron comprising dendritically linked amino acid units I

in which  $\mathsf{R}^1$  is  $\mathsf{C}_{1.6}$ -alkylene and X is selected from the group consisting of -O-, -S-, -NH- and -CO-, and each unit of a dendron may have the same groups  $\mathsf{R}^1$  and X, and in which a first dendron has n (where n is 2) levels of dendritically linked amino acid units and  $\mathsf{2}^n$  terminal branches, to p (where p is at least 2) of which terminal branches there are linked anchor groups

where Y is selected from -CO-, -NH-, -O- and -S-, provided that at least one of X and Y is -CO-,

 $\rm H^2$  is an organic group containing at least one  $\rm C_{6-24}$ -alkyl, -alkenyl, or -alkynyl group,  $\rm H^3$  is selected from the group consisting of hydrogen, amine, blocked amine, hydroxyl,  $\rm C_{1-24}$  alkoxy, thiol, COOH, or an organic group containing at least one  $\rm C_{6-24}$ -alkyl, alkenyl or -alkynyl group,  $\rm C_{1-6}$ -alkanoyloxy, or  $\rm C_{1-6}$ -alkanamido,

and in which a second dendron has m (where m is in the range 3 to 5) levels of dendritically linked amino acid units of the formula I above in which the groups R¹ and X may be the same as or different to one another and the same as or different to those of the amino acid units in the first dendron and 2<sup>m</sup> terminal branches, each of which is either unconjugated and is a group selected from NH<sub>2</sub>, N\*H<sub>2</sub>R¹¹, in which R¹¹ is hydrogen or C<sub>1-4</sub> alkyI, COOH, COO¹, OH or SH, or is conjugated via the terminal -X-,-NH-or-CO- group to a group R¹² where R¹² is a methylol group, an active ligand or an organic group comprising a sugar moiety.

2. A compound according to claim 1 in which the focal group is an amino acid unit, having the formula I

in which X and R1 are as defined above.

- 3. A compound according to claim 2 in which X in the focal group is -CO- or -NH-.
- 4. A compound according to claim 3 in which X in the focal group is -NH-.
- A compound according to claim 4 in which the focal group is formed from lysine or ornithine, that is R<sup>1</sup> is (CH<sub>2</sub>)<sub>4</sub> or (CH<sub>2</sub>)<sub>3</sub>.
- 6. A compound according to any preceding claim in which  $2^n = p$ .
- 7. A compound according to any preceding claim in which each terminal branch of the second dendron is -NH2 or

-N+H2R11.

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- A compound according to any preceding claim in which in each of the groups of the formula I, the groups R<sup>1</sup> and X are the same.
- 9. A compound according to claim 8 in which X is -NH- and R1 is -(CH<sub>2</sub>)<sub>4</sub> or -(CH<sub>2</sub>)<sub>3</sub>-.
- 10. A compound according to any preceding claim which is bound to a resin support through the focal unit of the core.
- 10 11. A compound according to claim 10 which is bound to the support via a spacer.
  - 12. A composition comprising a compound of any preceding claim.
- 13. A pharmaceutical composition comprising a pharmaceutical excipient and a compound according to any of claims1 to 9.
  - 14. A method of synthesis in which a focal reagent which has two reactive groups is reacted in a first series of first dendron producing steps as follows:
    - 1. an amino acid reagent of the formula II

R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>16</sup>

COR<sup>15</sup>

II

in which R1 and X are as defined in claim 1

R14 is H when X is -O-, -S- or -NH-,

OH when X is -CO-, or

is a protecting group,

R15 is a carboxylic acid protecting group, hydroxyl, or a carboxylic acid activating group,

R16 is H, an amine protecting group or an amine activating group,

provided that at least two of  $R^{14}$ ,  $R^{15}$  and  $R^{16}$  is other than an activating group and at least one of  $R^{14}$ ,  $R^{15}$  and  $R^{16}$  is other than a protecting group, is reacted

with the focal reagent, optionally after a step in which the desired reactive group of the focal reagent is deprotected and/or activated whereby the reactive group on the focal reagent reacts with one of the groups R14X-, R15-CO- and R16-NH-;

 a second step in which both unreacted groups R<sup>14</sup>X-, R<sup>15</sup>CO- and R<sup>16</sup>NH- of the product of the preceding step are, if necessary, deprotected and/or activated and reacted with at least two equivalents of a trifunctional reagent having the general formula II,

R<sup>16</sup>XR<sup>1</sup>CH-NHR<sup>16</sup>
| II

in which the groups  $\mathbb{R}^1$ ,  $\mathbb{R}^{14}$ ,  $\mathbb{R}^{15}$  and  $\mathbb{R}^{16}$  are as defined in step 1 and are the same or different as in the trifunctional reagent used in step 1;

- 3. One repeat of step 2, using at least four equivalents of trifunctional reagent of the general formula II; and 4. An anchor group attachment step in which at least two of the four groups R14X-, R15CO- and R16NH- are,
- if necessary, deprotected and/or activated, and reacted with a lipophilic reagent of the formula III

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in which Y, R<sup>2</sup> and R<sup>3</sup> are as defined above, and R<sup>17</sup> is OH or a carboxylic acid activating group, where Y is -CO- or R<sup>17</sup> is H or an amine, hydroxyl or thiol activating group, respectively, where Y is -NH-, -O- or -S-, whereby the said at least two groups react with R<sup>17</sup>Y- to conjugate

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groups to the X, CO- or NH-; and

a second dendron forming series of reaction steps in which

1. the other of the reactive groups of the focal reagent is, in a step separate to step 1 of the first series of first dendron producing steps,

reacted with an amino acid reagent of the formula II

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in which R1 and X are as defined above,

R14 is H when X is -O-, -S- or -NH-,

OH when X is -CO-, or

is a protecting group,

R15 is a carboxylic acid protecting group, hydroxyl, or a carboxylic acid activating group,

R16 is H, an amine protecting group or an amine activating group,

provided that at least two of R<sup>14</sup>, R<sup>15</sup> and R<sup>16</sup> is other than an activating group and at least one of R<sup>14</sup>, R<sup>15</sup> and R<sup>16</sup> is other than a protecting group,

optionally after a step in which the desired reactive group of the focal reagent and/or one of the groups -XR<sup>14</sup>, -COR<sup>15</sup> and -NHR<sup>16</sup> is deprotected and/or activated whereby the other of the reactive groups on the focal reagent reacts with one of the groups R<sup>14</sup>X-, R<sup>15</sup>-CO- and R<sup>16</sup>-NH-;

2. a second step in which both unreacted groups R14X-, R15CO- and R16NH- of the product of the preceding step are, if necessary, deprotected and/or activated and reacted with at least two equivalents of a trifunctional reagent having the general formula II.

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in which the groups R1, R14, R15 and R16 are as defined in step 1 and are the same or different as in the trifunctional reagent used in step 1;

3. (m-1) repeats of step 2, using in each case at least 2 <sup>(r+1)</sup> equivalents of trifunctional reagent for the r<sup>th</sup> repeat of step 2, until m levels of dendritically linked amino acids have been formed, where m is in the range 3 to 5.

15. A method according to claim 14 involving a preliminary step of reacting a focal reagent of the formula VI

VI

in which R<sup>17</sup>, R<sup>18</sup> and R<sup>19</sup> are selected from the same groups as R<sup>14</sup>, R<sup>16</sup> and R<sup>15</sup>, respectively, as defined in claim 14, with a substrate having a pendant group which is capable of reacting with one of the groups -XR<sup>17</sup>, -NHR<sup>18</sup> and -COR<sup>19</sup>, optionally after deprotection and/or activation of the said pendant group, whereby the focal reagent is bound to the substrate.

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16. A method according to claim 15 in which the substrate is an immobile support, preferably a resin, more preferably a polyacrylamide-based resin.

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17. A method according to claim 16 in which the resin has pendant amine groups and in which the group -COR<sup>19</sup> is reacted with said pendant amine groups in the presence of an activating compound to form a peptide bond.

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18. A method according to claim 17 in which R<sup>17</sup> and R<sup>18</sup> are each different amine protecting groups.

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19. A method according to any of claims 14 to 18 in which in each of the steps in each respective series the reagent of the formula II is the same, preferably in which the reagent of the formula II is the same for each series.

20. A method according to claim 19 in which X is -NH- and in which the groups R<sup>14</sup> and R<sup>15</sup> are the same amino protecting groups.

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#### Patentansprüche

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 Dendritische Verbindung, umfassend einen Kem, der eine fokale Gruppe einschließt, von der mindestens zwei Dendrons ausgehen, wobei jedes Dendron dendritisch verknüpfte Aminosäureeinheiten I

> X-R¹-CH-NH |

co-

umfaßt, worin  $\mathbb{R}^1$   $\mathbb{C}_{1.6}$ -Alkylen ist und X aus der Gruppe, bestehend aus -O-, -S-, -NH- und -CO-, ausgewählt ist und jede Einheit eines Dendrons die gleichen Gruppen  $\mathbb{R}^1$  und X aufweisen kann, und worin ein erstes Dendron n (wobei n 2 ist) Ebenen von dendritisch verknüpften Aminosäureeinheiten und  $2^n$  endständige Zweige aufweist, wobei mit p (wobei p mindestens 2 ist) endständigen Zweigen Ankergruppen

-Y-CH-R<sup>3</sup> | |R<sup>2</sup>

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verknüpft sind, worin Y aus -CO-, -NH-, -O- und -S- ausgewählt ist, mit der Maßgabe, daß mindestens eines von X und Y -CO- ist,

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 $R^2$  eine organische Gruppe ist, die mindestens eine  $C_{6\cdot 24}$ -Alkyl-, -Alkenyl- oder -Alkinylgruppe enthält,  $R^3$  aus der Gruppe, bestehend aus Wasserstoff, Amin, blockiertem Amin, Hydroxyl,  $C_{1\cdot 24}$ -Alkoxy, Thiol, COOH oder einer organischen Gruppe, die mindestens eine  $C_{6\cdot 24}$ -Alkyl, -Alkenyl oder -Alkinylgruppe,  $C_{1\cdot 6}$ -Alkanoyloxy oder  $C_{1-6}$ -Alkanamido enthält, ausgewählt ist,

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und worin ein zweites Dendron m (wobei m im Bereich von 3 bis 5 liegt) Ebenen von dendritisch verknüpften Aminosäureeinheiten der Formel I oben, worin die Gruppen  $\mathbb{R}^1$  und X gleich oder verschieden voneinander und gleich denjenigen der Aminosäureeinheiten im ersten Dendron oder davon verschieden sein können, und  $2^m$  end-

ständige Zweige aufweist, von denen ein jeder entweder unkonjugiert ist und eine Gruppe darstellt, die aus NH<sub>2</sub>, N+H<sub>2</sub>R<sup>11</sup>, worin R<sup>11</sup> Wasserstoff oder C<sub>1-4</sub>-Alkyl, COOH, COOH

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2. Verbindung nach Anspruch 1, worin die fokale Gruppe eine Aminosäureeinheit der Formel I

-X-R1- CH -NH-| | CO-

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ist, worin X und R1 wie oben definiert sind.

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- 3. Verbindung nach Anspruch 2, worin X in der fokalen Gruppe -CO- oder -NH- ist.
- 4. Verbindung nach Anspruch 3, worin X in der fokalen Gruppe -NH- ist.

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- Verbindung nach Anspruch 4, worin die fokale Gruppe von Lysin oder Omithin gebildet wird, d.h., R¹ ist -(CH<sub>2</sub>)<sub>4</sub>oder -(CH<sub>2</sub>)<sub>3</sub>-.
- 6. Verbindung nach irgendeinem der vorhergehenden Ansprüche, worin 2<sup>n</sup> = p ist.

 Verbindung nach irgendeinem der vorhergehenden Ansprüche, worin jeder endständige Zweig des zweiten Dendrons -NH<sub>2</sub> oder -N+H<sub>2</sub>R<sup>11</sup> ist.

 Verbindung nach irgendeinem der vorhergehenden Ansprüche, worin die Gruppen R¹ und X in jeder der Gruppen der Formel I gleich sind.

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9. Verbindung nach Anspruch 8, worin X -NH- ist und R1 -(CH<sub>2</sub>)<sub>4</sub>- oder -(CH<sub>2</sub>)<sub>3</sub>- ist.

 Verbindung nach irgendeinem der vorhergehenden Ansprüche, welche über die fokale Einheit des Kerns an einen Harzträger gebunden ist.

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- 11. Verbindung nach Anspruch 10, welche über einen Spacer an den Träger gebunden ist.
- 12. Zusammensetzung, umfassend eine Verbindung nach irgendeinem der vorhergehenden Ansprüche.

40 13. Pharmazeutische Zusammensetzung, umfassend einen pharmazeutischen Exzipienten und eine Verbindung nach irgendeinem der Ansprüche 1 bis 9.

14. Syntheseverfahren, worin ein fokales Reagenz, welches zwei reaktive Gruppen aufweist, umgesetzt wird in einer ersten Folge von Schritten zur Bildung eines ersten Dendrons wie folgt:

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1. ein Aminosäurereagenz der Formel II

R¹4XR¹CH-NHR¹6

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worin R1 und X wie in Anspruch 1 definiert sind,

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R14 H ist, wenn X -O-, -S- oder -NH- ist,

OH ist, wenn X -CO- ist, oder eine Schutzgruppe ist,

R15 eine Carbonsäure-Schutzgruppe, Hydroxyl oder eine Carbonsäure-aktivierende Gruppe ist,

 $R^{16}$  H, eine Amin-Schutzgruppe oder eine Amin-aktivierende Gruppe ist, vorausgesetzt, daß mindestens zwei von  $R^{14}$ ,  $R^{15}$  und  $R^{16}$  keine aktivierende Gruppe sind und mindestens eines von  $R^{14}$ ,  $R^{15}$  und  $R^{16}$  keine Schutzgruppe ist,

wird mit dem fokalen Reagenz umgesetzt, gegebenenfalls nach einem Schritt, in dem die gewünschte reaktive Gruppe des fokalen Reagenz von der Schutzgruppe befreit und/oder aktiviert wird, wodurch die reaktive Gruppe auf dem fokalen Reagenz mit einer der Gruppen R<sup>14</sup>X-, R<sup>15</sup>CO- und R<sup>16</sup>NH- reagiert;

 ein zweiter Schritt, in dem beide nicht umgesetzten Gruppen R<sup>14</sup>X-, R<sup>15</sup>CO- und R<sup>16</sup>NH- des Produkts des vorhergehenden Schritts erforderlichenfalls von der Schutzgruppe befreit und/oder aktiviert werden und mit mindestens zwei Äquivalenten eines trifunktionellen Reagenz der allgemeinen Formel II

> R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>15</sup> | II | COR<sup>15</sup>

umgesetzt werden, worin die Gruppen R<sup>1</sup>, R<sup>14</sup>, R<sup>15</sup> und R<sup>16</sup> wie in Schritt 1 definiert sind und die gleichen wie in dem in Schritt 1 eingesetzten trifunktionellen Reagenz oder davon verschieden sind;

- 3. eine Wiederholung von Schritt 2, wobei mindestens vier Äquivalente des trifunktionellen Reagenz der allgemeinen Formel II eingesetzt werden; und
- 4. ein Ankergruppen-Verknüpfungsschritt, worin mindestens zwei der vier Gruppen R<sup>14</sup>X-, R<sup>15</sup>CO- und R<sup>16</sup>NHerforderlichenfalls von der Schutzgruppe befreit und/oder aktiviert werden und mit einem lipophilen Reagenz der Formei III

R'Y-CH-R³ | III | R²

umgesetzt werden,

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worin Y,  $R^2$  und  $R^3$  wie oben definiert sind und  $R^{17}$  OH oder eine Carbonsäure-aktivierende Gruppe ist, wenn Y -CO- ist.

oder R<sup>17</sup> H oder eine Amin-, Hydroxyl- oder Thiol-aktivierende Gruppe ist, wenn Y -NH-, -O- oder -S- ist, wodurch die genannten mindestens zwei Gruppen mit R<sup>17</sup>Y- reagieren, um

-Y-CH-R³-Gruppen | | | R²

mit dem X, CO- oder NH- zu konjugieren;

und einer Folge von Reaktionsschritten zur Bildung eines zweiten Dendrons, beinhaltend

1. daß die andere der reaktiven Gruppen des fokalen Reagenz in einem Schritt, welcher von Schritt 1 der ersten Folge von Schritten zur Bildung des ersten Dendrons getrennt ist, mit einem Aminosäurereagenz der Formel II

umgesetzt wird, worin R1 und X wie oben definiert sind,

R14 Hist, wenn X -O-, -S- oder -NH- ist.

OH ist, wenn X -CO- ist, oder eine Schutzgruppe ist,

R15 eine Carbonsäure-Schutzgruppe, Hydroxyl oder eine Carbonsäure-aktivierende Gruppe ist,

R16 H, eine Amin-Schutzgruppe oder eine Amin-aktivierende Gruppe ist,

vorausgesetzt, daß mindestens zwei von  $R^{14}$ ,  $R^{15}$  und  $R^{16}$  keine Aktivierungsgruppe sind und mindestens eines von  $R^{14}$ ,  $R^{15}$  und  $R^{16}$  keine Schutzgruppe ist;

gegebenenfalls nach einem Schritt, in dem die gewünschte reaktive Gruppe des fokalen Reagenz und/ oder eine der Gruppen -XR14, -COR15

und -NHR<sup>16</sup> von der Schutzgruppe befreit und/oder aktiviert wird, wodurch die andere der reaktiven Gruppen auf dem fokalen Reagenz mit einer der Gruppen R<sup>14</sup> X-, R<sup>15</sup>CO- und R<sup>16</sup>NH- reagiert;

2. einen zweiten Schritt, in dem beide nicht umgesetzten Gruppen R¹⁴X-, R¹⁵CO- und R¹⁶NH- des Produkts des vorhergehenden Schritts erforderlichenfalls von der Schutzgruppe befreit und/oder aktiviert werden und mit mindestens zwei Äquivalenten eines trifunktionellen Reagenz der allgemeinen Formel II

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R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>16</sup>

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umgesetzt werden, worin die Gruppen R<sup>1</sup>, R<sup>14</sup>, R<sup>15</sup> und R<sup>16</sup> wie in Schritt 1 definiert sind und die gleichen wie in dem bei Schritt 1 eingesetzten trifunktionellen Reagenz oder davon verschieden sind;

3. (m-1) Wiederholungen von Schritt 2, wobei in jedem Fall mindestens 2(r+1) Äquivalente des trifunktionellen Reagenz für die r. Wieder-holung des Schritts 2 eingesetzt werden, bis m Ebenen von dendritisch verknüpften Aminosäuren gebildet wurden, wobei m im Bereich von 3 bis 5 liegt.

 Verfahren nach Anspruch 14, beinhaltend einen einleitenden Schritt der Umsetzung eines fokalen Reagenz der Formel VI

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R<sup>17</sup>-X-R<sup>1</sup>-CH-NH-R<sup>1</sup>

VI

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worin R<sup>17</sup>, R<sup>18</sup> und R<sup>19</sup> aus den gleichen Gruppen wie R<sup>14</sup>, R<sup>16</sup> bzw. R<sup>15</sup> wie in Anspruch 14 definiert ausgewählt sind, mit einem Substrat, das eine Seitengruppe aufweist, welche zur Reaktion mit einer der Gruppen -XR<sup>17</sup>, -NHR<sup>18</sup> und -COR<sup>19</sup> in der Lage ist, gegebenenfalls nach Befreiung von der Schutzgruppe und/oder Aktivierung der Seitengruppe, wodurch das fokale Reagenz an das Substrat gebunden wird.

- Verfahren nach Anspruch 15, worin das Substrat ein immobiler Träger, vorzugsweise ein Harz, noch bevorzugter ein Harz auf Polyacrylamidbasis, ist.
- 17. Verfahren nach Anspruch 16, worin das Harz seitenständige Amingruppen aufweist und worin die Gruppe -COR19 mit diesen seitenständigen Amingruppen in Gegenwart einer aktivierenden Verbindung umgesetzt wird, um eine Peptidbindung zu bilden.
  - 18. Verfahren nach Anspruch 17, worin R<sup>17</sup> und R<sup>18</sup> jeweils unterschiedliche Amin-Schutzgruppen sind.

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- 19. Verfahren nach irgendeinem der Ansprüche 14 bis 18, worin in jedem der Schritte in einer der jeweiligen Folgen das Reagenz der Formel II gleich ist, vorzugsweise, worin das Reagenz der Formel II für jede Folge das gleiche ist.
- 20. Verfahren nach Anspruch 19, worin X -NH- ist und worin die Gruppen R14 und R15 die gleichen Amino-Schutzgruppen sind.

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#### Revendications

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 Composé dendritique comprenant un noyau incluant un groupe focal d'où s'étendent au moins deux dendrons, chaque dendron comprenant des unités d'aminoacide | liées de manière dendritique

où R¹ est alkylène en C<sub>1-6</sub> et X est choisi dans le groupe consistant en -O-, -S-, -NH- et -CO-, et chaque unité d'un dendron peut avoir les mêmes groupes R¹ et X, et où un premier dendron a n (où n est 2) niveaux d'unités d'aminoacide liées de manière dendritique et 2<sup>n</sup> branches terminales, branches terminales à p desquelles (où p est au moins 2) sont liés des groupes d'ancrage

où Y est choisi parmi -CO-, -NH-, -O- et -S-, à condition qu'au moins l'un des groupes X et Y soit -CO-,

 $R^2$  est un groupe organique contenant au moins un groupe alkyle, alcényle ou alcynyle en  $C_{6\cdot 24}$ ,  $R^3$  est choisi dans le groupe consistant en l'hydrogène, amine, amine bloquée, hydroxyle, alcoxy en  $C_{1\cdot 24}$ , thiol, COOH ou un groupe organique contenant au moins un groupe alkyle, alcényle ou alcynyle en  $C_{6\cdot 24}$ , alcanoyloxy en  $C_{1\cdot 6}$  ou alcanamido en  $C_{1\cdot 6}$ .

et où un second dendron a m (où m est dans le domaine de 3 à 5) niveaux d'unités d'aminoacide liées de manière dendritique de formule I ci-dessus où les groupes R¹ et X peuvent être identiques ou différents l'un de l'autre et identiques ou différents de ceux des unités d'aminoacide dans le premier dendron et 2<sup>m</sup> branches terminales, dont chacune est non conjuguée et est un groupe choisi parmi NH<sub>2</sub>, N\*H<sub>2</sub>R¹¹ où R¹¹ est l'hydrogène ou alkyle en C<sub>1.4</sub>, COOH, COO¹, OH ou SH, ou est conjuguée par le groupe -X-, -NH- ou -CO- terminal à un groupe R¹² où R¹² est un groupe méthylol, un ligand actif ou un groupe organique comprenant une entité glucidique.

2. Composé selon la revendication 1, où le groupe focal est une unité d'aminoacide ayant la formule l

où X et R1 sont définis comme ci-dessus.

- 3. Composé selon la revendication 2, où X dans le groupe focal est -CO- ou -NH-.
- 4. Composé selon la revendication 3, où X dans le groupe focal est -NH-.
- Composé selon la revendication 4, où le groupe focal est constitué par de la lysine ou de l'ornithine, c'est-à-dire que R¹ est (CH<sub>2</sub>)<sub>4</sub> ou (CH<sub>2</sub>)<sub>3</sub>.
  - 6. Composé selon l'une quelconque des revendications précédentes, où  $2^n = p$ .
- Composé selon l'une quelconque des revendications précédentes, où chaque branche terminale du second dendron est -NH<sub>2</sub> ou -N+H<sub>2</sub>R<sup>11</sup>.
  - 8. Composé selon l'une quelconque des revendications précédentes, où, dans chacun des groupes de formule I, les

groupes R1 et X sont les mêmes.

- 9. Composé selon la revendication 8, où X est -NH- et R1 est -(CH<sub>2</sub>)<sub>4</sub> ou -(CH<sub>2</sub>)<sub>3</sub>-.
- Composé selon l'une quelconque des revendications précédentes, qui est lié à un support résinique par l'unité focale du noyau.
  - 11. Composé selon la revendication 10, qui est lié au support par le biais d'un espaceur.
- 10 12. Composition comprenant un composé selon l'une quelconque des revendications précédentes.
  - Composition pharmaceutique comprenant un excipient pharmaceutique et un composé selon l'une quelconque des revendications 1 à 9.
- 15 14. Procédé de synthèse où un réactif local qui a deux groupes réactifs est mis à réagir dans une première série d'étapes de production d'un premier dendron de la manière suivante :
  - 1. un réactif aminoacide de formule II

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## R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>16</sup> COR<sup>15</sup>

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où R1 et X sont définis comme dans la revendication 1

R14 est H quand X est -O-, -S- ou -NH-,

OH quand X est -CO-, ou

est un groupe protecteur.

 $\mathsf{R}^{15}$  est un groupe protecteur d'acide carboxylique, un groupe hydroxyle ou un groupe activateur d'acide carboxylique,

R16 est H, un groupe protecteur d'amine ou un groupe activateur d'amine,

à condition qu'au moins deux des groupes  $R^{14}$ ,  $R^{15}$  et  $R^{16}$  ne soient pas un groupe activateur et qu'au moins l'un des groupes  $R^{14}$ ,  $R^{15}$  et  $R^{16}$  ne soit pas un groupe protecteur,

est mis à réagir avec le réactif focal, éventuellement après une étape au cours de laquelle le groupe réactif souhaité du réactif focal est déprotégé et/ou activé de sorte que le groupe réactif sur le réactif focal réagit avec l'un des groupes R<sup>14</sup>X-, R<sup>15</sup>-CO- et R<sup>16</sup>-NH-;

2. une seconde étape au cours de laquelle les deux groupes R<sup>14</sup>X-, R<sup>15</sup>-CO- et R<sup>16</sup>NH- qui n'ont pas réagi du produit de l'étape précédente sont, si nécessaire, déprotégés et/ou activés et mis à réagir avec au moins deux équivalents d'un réactif trifonctionnel ayant la formule générale II.

R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>16</sup> COR<sup>15</sup>

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où les groupes R1, R14, R15 et R16 sont définis comme dans l'étape 1 et sont identiques ou différents de ceux du réactif trifonctionnel utilisé dans l'étape 1 ;

3. une répétition de l'étape 2 au moyen d'au moins 4 équivalents de réactif trifonctionnel de formule générale II ; et

4. une étape de fixation de groupes d'ancrage au cours de laquelle au moins deux des quatre groupes R¹4X., R¹5CO- et R¹6NH- sont, si nécessaire, déprotégés et/ou activés, et mis à réagir avec un réactif lipophile de formule III

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où Y, R<sup>2</sup> et R<sup>3</sup> sont définis comme ci-dessus, et R<sup>17</sup> est OH ou un groupe activateur d'acide carboxylique quand Y est -CO-, ou bien R17 est H ou un groupe activateur d'amine, d'hydroxyle ou de thiol, respectivement, quand Y est -NH-, -O- ou -S-,

de sorte que lesdits groupes au nombre d'au moins deux réagissent avec R17Y-pour conjuguer des groupes

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à X, CO- ou NH- ; et

une seconde série d'étapes réactionnelles de formation d'un second dendron où

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1. dans une étape séparée de l'étape 1 de la première série d'étapes de production d'un premier dendron, l'autre des groupes réactifs du réactif focal est mis à réagir avec un réactif aminoacide de formule II

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II

où R1 et X sont définis comme ci-dessus.

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R14 est H quand X est -O-, -S- ou -NH-,

OH quand X est -CO-, ou

est un groupe protecteur,

R15 est un groupe protecteur d'acide carboxylique, un groupe hydroxyle ou un groupe activateur d'acide

R16 est H, un groupe protecteur d'amine ou un groupe activateur d'amine,

à condition qu'au moins deux des groupes R14, R15 et R16 ne soient pas un groupe activateur et qu'au moins t'un des groupes R14, R15 et R16 ne soit pas un groupe protecteur,

éventuellement après une étape au cours de laquelle le groupe réactif souhaité du réactif focal et/ou l'un des groupes -XR14, -COR15 et -NHR16 est déprotégé et/ou activé de sorte que l'autre des groupes réactifs du réactif focal réagit avec l'un des groupes R14X-, R15-CO- et R16-NH- ;

2. une seconde étape au cours de laquelle les deux groupes R14X-, R15CO- et R16NH- qui n'ont pas réagi du produit de l'étape précédente sont, si nécessaire, déprotégés et/ou activés et mis à réagir avec au moins deux équivalents d'un réactif trifonctionnel ayant la formule générale II,

R<sup>14</sup>XR<sup>1</sup>CH-NHR<sup>16</sup> COR<sup>15</sup>

П

où les groupes R1, R14, R15 et R16 sont définis comme dans l'étape 1 et sont identiques ou différents de ceux du réactif trifonctionnel utilisé dans l'étape 1 ;

3. (m-1) répétitions de l'étape 2, utilisant dans chaque cas au moins 2<sup>(r+1)</sup> équivalents de réactif trifonctionnel pour la r<sup>ème</sup> répétition de l'étape 2, jusqu'à ce que m<sub>i</sub>niveaux d'aminoacides liés de manière dendritique aient été formés, où m est dans le domaine de 3 à 5.

15. Procédé selon la revendication 14, mettant en jeu une étape préliminaire de réaction d'un réactif focal de formule VI

R17-X-R1-CH-NH-R18

CO R19

VI

où R<sup>17</sup>, R<sup>18</sup> et R<sup>19</sup> sont choisis dans les mêmes groupes que R<sup>14</sup>, R<sup>16</sup> et R<sup>15</sup>, respectivement, tels que définis dans la revendication 14, avec un substrat ayant un groupe latéral qui est capable de réagir avec l'un des groupes -XR<sup>17</sup>, -NHR<sup>18</sup> et -COR<sup>19</sup>, éventuellement après déprotection et/ou activation dudit groupe latéral, de sorte que le réactif focal est lié au substrat.

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- 16. Procédé selon la revendication 15, où le substrat est un support immobile, de préférence une résine, de préférence encore une résine à base de polyacrylamide.
- 17. Procédé selon la revendication 16, où la résine a des groupes amine latéraux et où le groupe -COR19 est mis à réagir avec lesdits groupes amine latéraux en présence d'un composé activateur pour former une liaison peptidique.
- 20 18. Procédé selon la revendication 17, où R<sup>17</sup> et R<sup>18</sup> sont chacun des groupes protecteurs d'amine différents.
  - 19. Procédé selon l'une quelconque des revendications 14 à 18 où, dans chacune des étapes dans chaque série respective, le réactif de formule It est le même, de préférence où le réactif de formule It est le même pour chaque série.
  - 20. Procédé selon la revendication 19, où X est -NH- et où les groupes R<sup>14</sup> et R<sup>15</sup> sont les mêmes groupes protecteurs d'amino.

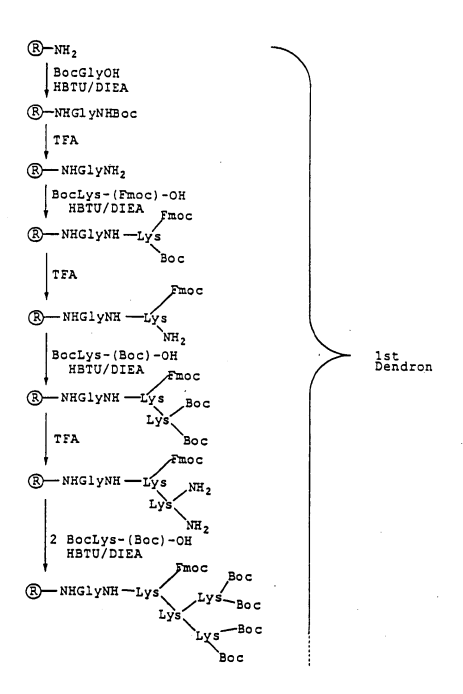


Figure 1.

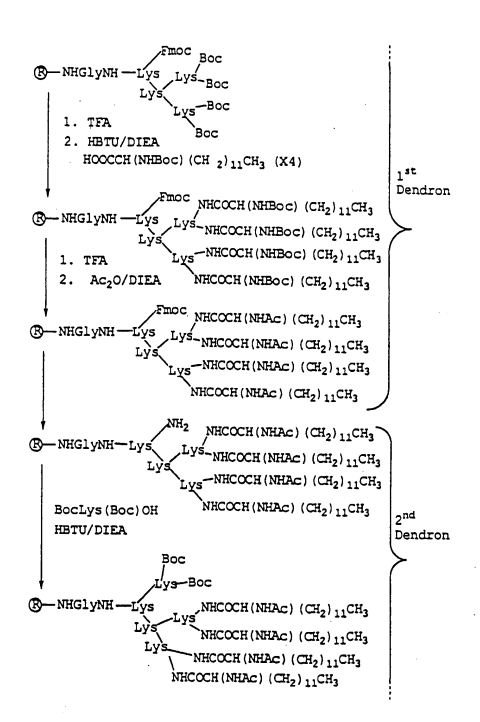
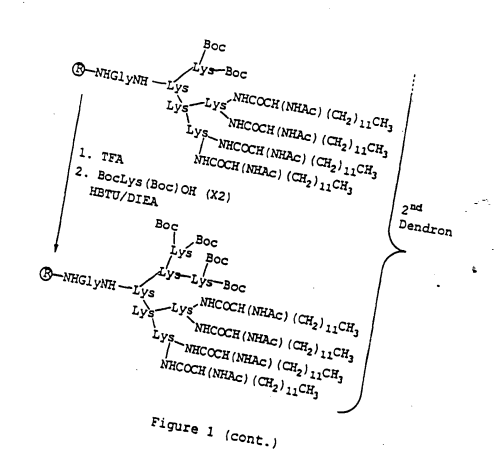
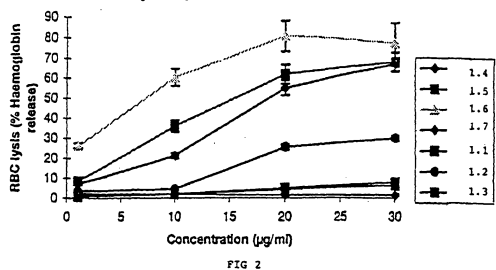


Figure 1 (cont.)



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# Haemoglobin release after th. Incubation of rat erythrocytes with dendrimers



# Relationship between the number of amino groups and haemoglobin release at 20 µg/ml after 1h. incubation

